Storypoint Prediction - titanium

September 6, 2024

1 Storypoint Prediction: Regression Approach

1.1 Preparation

```
[]: import os
     import json
     import matplotlib.pyplot as plt
     import numpy as np
     import pandas as pd
     import seaborn as sns
     from scipy.sparse import csr_matrix, hstack, vstack
     from sklearn.pipeline import Pipeline
     from sklearn.preprocessing import RobustScaler
     from sklearn.metrics import mean_squared_error, r2_score, mean_absolute_error, u

¬f1_score, precision_score, recall_score, accuracy_score
     from sklearn.feature_extraction.text import CountVectorizer
     from sklearn.model_selection import cross_val_score, learning_curve, u
      ⇒validation curve
     from trainer import GridSearchCVTrainer
     project_name = 'titanium'
```

1.1.1 Plot learning curve

```
# Generate learning curve data
  train_sizes, train_scores, test_scores = learning_curve(
       estimator, X, y, cv=cv, n_jobs=n_jobs, train_sizes=train_sizes,__
⇔scoring='neg_mean_squared_error')
  train scores mean = np.mean(train scores, axis=1) # Calculate mean of |
⇔training scores
  train_scores_std = np.std(train_scores, axis=1) # Calculate standard_
→ deviation of training scores
  test scores mean = np.mean(test scores, axis=1) # Calculate mean of test
\hookrightarrowscores
  test_scores_std = np.std(test_scores, axis=1) # Calculate standard_
→deviation of test scores
  plt.grid() # Display grid
  # Fill the area between the mean training score and the mean +/- stdu
⇔training score
  plt.fill_between(train_sizes, train_scores_mean - train_scores_std,
                    train_scores_mean + train_scores_std, alpha=0.1,
                    color="r")
  # Fill the area between the mean test score and the mean +/- std test score
  plt.fill_between(train_sizes, test_scores_mean - test_scores_std,
                   test_scores_mean + test_scores_std, alpha=0.1, color="g")
  # Plot mean training score as points
  plt.plot(train sizes, train scores mean, 'o-', color="r",
           label="Training score")
  # Plot mean test score as points
  plt.plot(train_sizes, test_scores_mean, 'o-', color="g",
           label="Validation score")
  plt.legend(loc="best") # Display legend
  return plt
```

1.1.2 Plot validation curve

```
# Calculate mean and standard deviation of training and validation scores
  train_mean = np.mean(train_scores, axis=1)
  tran_std = np.std(train_scores, axis=1)
  val_mean = np.mean(val_scores, axis=1)
  val_std = np.std(val_scores, axis=1)
  print(val_mean)
  # Plot train scores
  plt.plot(param_range, train_mean, color='r', marker='o', markersize=5,u
⇔label='Training score')
  plt.fill_between(param_range, train_mean + tran_std, train_mean - tran_std,__
⇔alpha=0.15, color='r')
  # Plot validation scores
  plt.plot(param_range, val_mean, color='g', linestyle='--', marker='s',u
→markersize=5, label='Validation score')
  plt.fill_between(param_range, val_mean + val_std, val_mean - val_std,__
⇒alpha=0.15, color='g')
  plt.title(title) # Set title of the plot
  plt.grid()
                         # Display grid
  plt.xscale('log')
                        # Set x-axis scale to log
  plt.legend(loc='best') # Display legend
  plt.xlabel('Parameter') # Set x-axis label
  plt.ylabel('Score')
                         # Set y-axis label
  # Set y-axis limits
  if y_lim != None:
      plt.ylim(y_lim)
  return plt
```

1.1.3 Evaluate model

```
lines.append(f' - Mean squared error:
                                           {mse:.2f}')
  lines.append(f' - Root mean squared error: {rmse:.2f}')
  lines.append(f' - Mean absolute error:
                                           {mae:.2f}')
                                           {r2:.2f}')
  lines.append(f' - R2 error:
  y_pred = np.round(y_pred).astype(int)
  f1 = f1_score(y_test, y_pred, average='weighted')
  precision = precision_score(y_test, y_pred, average='weighted',__
→zero_division=0)
  recall = recall_score(y_test, y_pred, average='weighted', zero_division=0)
  accuracy = accuracy_score(y_test, y_pred)
                                           {f1:.2f}')
  lines.append(f' - F1 score:
  lines.append(f' - Precision:
                                           {precision:.2f}')
                                          {recall:.2f}')
  lines.append(f' - Recall:
  lines.append(f' - Accuracy:
                                           {accuracy:.2f}')
  lines.append('----')
  lines.append('')
  # Save to file
  if(save_directory != None):
      filename = save_directory + project_name + '.txt'
      directory = os.path.dirname(filename)
      if not os.path.exists(directory):
          os.makedirs(directory)
      with open(filename, 'a') as f:
          for line in lines:
             print(line)
             f.write(line + '\n')
  else:
      for line in lines:
         print(line)
```

1.1.4 Set random seed

```
[]: import numpy as np
import random
import os

# Set random seed for numpy
np.random.seed(42)

# Set random seed for random
random.seed(42)

# Set random seed for os
```

```
os.environ['PYTHONHASHSEED'] = '42'
```

1.2 Dataset set-up

1.2.1 Bag of Words preprocessing

This is a Bag of Words preprocess approach. I will use 2 CountVectorizer from sklearn to change title and description to two 2 vectors and then concatenate them together. In the rest of this notebook, I will use cross-validation instead hold-out. Therefore, I will join the validation set with training set.

NOTE: I don't add the length of description (and the title) because it has a big deviation which may cause noise.

```
[]: # Import and remove NaN value
     data train = pd.concat([pd.read csv('data/' + project name + '/' + project name<sub>||</sub>
      →+ '_train.csv'),
                             pd.read_csv('data/' + project_name + '/' + project_name_
      →+ '_valid.csv')])
     data_test = pd.read_csv('data/' + project_name + '/' + project_name + '_test.
      ⇔csv')
     data_train['description'].replace(np.nan, '', inplace=True)
     data_test['description'].replace(np.nan, '', inplace=True)
     # Vectorize title
     title_vectorizer = CountVectorizer(ngram_range=(1, 2), min_df=2)
     title vectorizer.fit(pd.concat([data train['title'], data test['title']]))
     # Vectorize description
     description_vectorizer = CountVectorizer(ngram_range=(1, 2), min_df=2)
     description_vectorizer.fit(pd.concat([data_train['description'],__

¬data_test['description']]))
     X train = hstack([title vectorizer.transform(data_train['title']).astype(float),
                       description_vectorizer.transform(data_train['description']).
      ⇒astype(float),
                        data_train['title'].apply(lambda x : len(x)).to_numpy().
      \rightarrowreshape(-1, 1),
                       data_train['description'].apply(lambda x : len(x)).to_numpy().
      \rightarrowreshape(-1, 1)
     y_train = data_train['storypoint'].to_numpy().astype(float)
```

```
X test = hstack([title vectorizer.transform(data_test['title']).astype(float),
                       description_vectorizer.transform(data_test['description']).
      →astype(float),
                       data_test['title'].apply(lambda x : len(x)).to_numpy().
      \rightarrowreshape(-1, 1),
                       data_test['description'].apply(lambda x : len(x)).to_numpy().
      \rightarrowreshape(-1, 1)
     y_test = data_test['storypoint'].to_numpy().astype(float)
[]: print('Check training dataset\'shape:', X_train.shape, y_train.shape)
     print('Check testing dataset\'shape:', X_test.shape, y_test.shape)
    Check training dataset'shape: (2026, 20875) (2026,)
    Check testing dataset'shape: (225, 20875) (225,)
    I will use log-scale the label to get a normal distribution of it.
[]: y_train_log = np.log(y_train)
    1.3 Model training
    1.3.1 Linear Regressor
[]: from sklearn.linear model import ElasticNet
    Define params-grid:
[]: dict_param = {
         'alpha': [.0001, .001, .01, .1, 1, 10, 100, 1000, 10000],
         'l1_ratio': [.0, .2, .4, .6, .8, 1],
         'max_iter': [3000],
         'random_state': [42]
[]:|gridsearch = GridSearchCVTrainer(name='Elastic Net', model=ElasticNet(),__
      →param_grid=dict_param, cv=5, n_jobs=5)
     gridsearch.load_if_exists()
     gridsearch.fit(X_train, y_train_log)
     elastic_model = gridsearch.best_estimator_
     elastic_model.fit(X_train, y_train_log)
    There is no checkpoint file for this model.
               | 54/54 [26:37<00:00, 29.59s/it]
```

[]: ElasticNet(alpha=0.01, l1_ratio=1, max_iter=3000, random_state=42)

```
→y_logscale=True, save_directory='results/BoW/')
    Elastic Net model's evaluation results:
     - Mean squared error:
                                 12.63
     - Root mean squared error: 3.55
     - Mean absolute error:
                                2.17
     - R2 error:
                                -0.02
     - F1 score:
                                0.20
     - Precision:
                                0.27
     - Recall:
                                0.19
                                0.19
     - Accuracy:
[]: elastic_model.get_params()
[]: {'alpha': 0.01,
      'copy_X': True,
      'fit_intercept': True,
      'l1_ratio': 1,
      'max_iter': 3000,
      'positive': False,
      'precompute': False,
      'random_state': 42,
      'selection': 'cyclic',
      'tol': 0.0001,
      'warm_start': False}
    1.3.2 Support Vector Regressor
[]: from sklearn.svm import SVR
[]: dict_param = {
         'C': [.0001, .001, .01, .1, 1, 10, 100, 1000, 10000],
         'gamma': np.logspace(-9, 3, 13),
         'kernel': ['rbf']
     }
[]:|grid_search = GridSearchCVTrainer(name="Support Vector Regressor",model=SVR(),__
      →param_grid=dict_param, cv=5, n_jobs=5)
     grid_search.load_if_exists()
     grid_search.fit(X_train, y_train_log)
     svr_model = grid_search.best_estimator_
     svr_model.fit(X_train, y_train_log)
    There is no checkpoint file for this model.
```

[]: evaluate_model(elastic_model, 'Elastic Net model', X_test, y_test, u

```
100%|
              | 117/117 [03:04<00:00, 1.58s/it]
[]: SVR(C=1000, gamma=1e-08)
[]: evaluate_model(svr_model, 'SVR model', X_test, y_test, y_logscale=True,__
      ⇔save_directory='results/BoW/')
    SVR model's evaluation results:
     - Mean squared error:
                                 12.58
     - Root mean squared error: 3.55
     - Mean absolute error:
                                2.19
     - R2 error:
                                -0.01
     - F1 score:
                                0.21
     - Precision:
                                0.16
     - Recall:
                                0.30
     - Accuracy:
                                0.30
[]: svr_model.get_params()
[]: {'C': 1000,
      'cache_size': 200,
      'coef0': 0.0,
      'degree': 3,
      'epsilon': 0.1,
      'gamma': 1e-08,
      'kernel': 'rbf',
      'max_iter': -1,
      'shrinking': True,
      'tol': 0.001,
      'verbose': False}
    1.3.3 Random Forest Regressor
[]: from sklearn.ensemble import RandomForestRegressor
[]: dict_param = {
         'max_depth' : [1000, 2000, 5000],
         'min_samples_split': [25, 200, 1000],
         'min_samples_leaf': [1, 2, 3, 4],
         'max_features': [50, 100, 200],
         'n_estimators': [1024],
         'random_state': [42]
     }
[]: grid_search = GridSearchCVTrainer(name="Random Forest Regressor",
                                       model=RandomForestRegressor(),
```

```
param_grid=dict_param, cv = 5, n_jobs=5)
     grid_search.load_if_exists()
     grid_search.fit(X_train, y_train_log)
     rfr_model = grid_search.best_estimator_
    rfr_model.fit(X_train, y_train_log)
    There is no checkpoint file for this model.
    100%|
               | 108/108 [15:49<00:00, 8.80s/it]
[]: RandomForestRegressor(max_depth=1000, max_features=200, min_samples_split=25,
                           n_estimators=1024, random_state=42)
[]: evaluate_model(rfr_model, 'Random Forest model', X_test, y_test, u
      →y_logscale=True, save_directory='results/BoW/')
    Random Forest model's evaluation results:
     - Mean squared error:
     - Root mean squared error: 3.49
     - Mean absolute error:
                                 2.00
     - R2 error:
                                 0.02
     - F1 score:
                                0.23
                                 0.27
     - Precision:
     - Recall:
                                 0.25
     - Accuracy:
                                 0.25
[]: rfr_model.get_params()
[]: {'bootstrap': True,
      'ccp_alpha': 0.0,
      'criterion': 'squared_error',
      'max_depth': 1000,
      'max features': 200,
      'max_leaf_nodes': None,
      'max_samples': None,
      'min_impurity_decrease': 0.0,
      'min_samples_leaf': 1,
      'min_samples_split': 25,
      'min_weight_fraction_leaf': 0.0,
      'monotonic_cst': None,
      'n_estimators': 1024,
      'n_jobs': None,
      'oob_score': False,
      'random_state': 42,
      'verbose': 0,
      'warm_start': False}
```

1.3.4 XGBoost

```
[]: from xgboost import XGBRegressor
[]: dict_param = {
         'eta': np.linspace(0.01, 0.2, 3),
         'gamma': np.logspace(-2, 2, 5),
         'max_depth': np.asarray([3, 5, 7, 9]).tolist(),
         'min_child_weight': np.logspace(-2, 2, 5),
         'subsample': np.asarray([0.5, .1]),
         'reg alpha': np.asarray([0.0, 0.05]),
         'n estimators': np.asarray([10, 20, 50, 100]).tolist(),
         'random state': [42]
    }
[]: grid_search = GridSearchCVTrainer(name='XGBoost_
     grid_search.load_if_exists()
    grid_search.fit(X_train, y_train_log)
    xgb_model = grid_search.best_estimator_
    xgb_model.fit(X_train, y_train_log)
    There is no checkpoint file for this model.
    100%|
              | 4800/4800 [1:00:31<00:00, 1.32it/s]
[]: XGBRegressor(base_score=None, booster=None, callbacks=None,
                 colsample_bylevel=None, colsample_bynode=None,
                 colsample_bytree=None, device=None, early_stopping_rounds=None,
                 enable categorical=False, eta=0.105, eval metric=None,
                 feature_types=None, gamma=0.01, grow_policy=None,
                 importance_type=None, interaction_constraints=None,
                 learning_rate=None, max_bin=None, max_cat_threshold=None,
                 max_cat_to_onehot=None, max_delta_step=None, max_depth=9,
                 max_leaves=None, min_child_weight=0.01, missing=nan,
                 monotone constraints=None, multi strategy=None, n estimators=20,
                 n_jobs=None, num_parallel_tree=None, ...)
[]: evaluate_model(xgb_model, 'XGBoost Regressor model', X_test, y_test,__
      →y_logscale=True, save_directory='results/BoW/')
    XGBoost Regressor model's evaluation results:
     - Mean squared error:
                               12.19
     - Root mean squared error: 3.49
     - Mean absolute error:
                               2.20
     - R2 error:
                               0.02
     - F1 score:
                               0.16
     - Precision:
                               0.31
```

```
- Recall: 0.12
- Accuracy: 0.12
```

[]: xgb_model.get_params()

```
[]: {'objective': 'reg:squarederror',
      'base_score': None,
      'booster': None,
      'callbacks': None,
      'colsample_bylevel': None,
      'colsample_bynode': None,
      'colsample_bytree': None,
      'device': None,
      'early_stopping_rounds': None,
      'enable_categorical': False,
      'eval_metric': None,
      'feature_types': None,
      'gamma': 0.01,
      'grow_policy': None,
      'importance_type': None,
      'interaction_constraints': None,
      'learning_rate': None,
      'max_bin': None,
      'max_cat_threshold': None,
      'max_cat_to_onehot': None,
      'max_delta_step': None,
      'max_depth': 9,
      'max_leaves': None,
      'min_child_weight': 0.01,
      'missing': nan,
      'monotone_constraints': None,
      'multi_strategy': None,
      'n estimators': 20,
      'n_jobs': None,
      'num_parallel_tree': None,
      'random_state': 42,
      'reg_alpha': 0.05,
      'reg_lambda': None,
      'sampling_method': None,
      'scale_pos_weight': None,
      'subsample': 0.5,
      'tree_method': None,
      'validate_parameters': None,
      'verbosity': None,
      'eta': 0.105}
```

1.3.5 LightGBM

```
[]: from lightgbm import LGBMRegressor
     from sklearn.model_selection import ParameterSampler
[]: | dict param = {
         'n_estimator': [10, 20, 50, 100, 200, 500],
         'max_depth': np.asarray([5, 7, 9, 11, 13]).tolist(),
         'num leaves': ((np.power(2, np.asarray([5, 7, 9, 11, 13])) - 1) * (0.55 +_{\sqcup}
      \hookrightarrow (0.65 - 0.55) * np.random.rand(5))).astype(int).tolist(),
         'min data in leaf': np.linspace(100, 1000, 4).astype(int).tolist(),
         'feature_fraction': np.linspace(0.6, 1, 3),
         'bagging_fraction': np.linspace(0.6, 1, 3),
         'learning_rate': [0.01],
         'verbose': [-1],
         'random_state': [42]
     }
     def custom_sampler(param_grid):
         for params in ParameterSampler(param_grid, n_iter=1e9):
             range_num_leaves = ((0.5 * (2**params['max_depth'] - 1)), (0.7 *_l)
      if(range_num_leaves[0] <= params['num_leaves'] <= range_num_leaves[1]):</pre>
                 for key, value in params.items():
                     params[key] = [value]
                 yield params
[]: grid_search = GridSearchCVTrainer(name='LightGBM Regressor', __
      →model=LGBMRegressor(),
                                     param grid=list(custom sampler(dict param)), cv___
     \Rightarrow= 5, n_jobs=2)
     grid_search.load_if_exists()
     grid_search.fit(X_train, y_train_log)
     lgbmr_model = grid_search.best_estimator_
     lgbmr_model.fit(X_train, y_train_log)
    c:\Users\aupho\AppData\Local\Programs\Python\Python311\Lib\site-
    packages\sklearn\model_selection\_search.py:320: UserWarning: The total space of
    parameters 5400 is smaller than n_iter=1000000000. Running 5400 iterations. For
    exhaustive searches, use GridSearchCV.
      warnings.warn(
    There is no checkpoint file for this model.
    100%|
               | 1080/1080 [08:00<00:00, 2.25it/s]
[]: LGBMRegressor(bagging_fraction=0.6, feature_fraction=1.0, learning_rate=0.01,
                   max_depth=11, min_data_in_leaf=100, n_estimator=10,
```

num_leaves=1248, random_state=42, verbose=-1)

```
[]: evaluate_model(lgbmr_model, 'LightGBM regressor model', X_test, y_test,__
      LightGBM regressor model's evaluation results:
     - Mean squared error:
     - Root mean squared error: 3.52
     - Mean absolute error:
                               2.15
     - R2 error:
                               0.00
     - F1 score:
                               0.20
     - Precision:
                               0.16
     - Recall:
                               0.25
                               0.25
     - Accuracy:
    c:\Users\aupho\AppData\Local\Programs\Python\Python311\Lib\site-
    packages\lightgbm\basic.py:1218: UserWarning: Converting data to scipy sparse
    matrix.
      _log_warning("Converting data to scipy sparse matrix.")
[]: lgbmr_model.get_params()
[]: {'boosting_type': 'gbdt',
      'class weight': None,
      'colsample_bytree': 1.0,
      'importance_type': 'split',
      'learning_rate': 0.01,
      'max depth': 11,
      'min_child_samples': 20,
      'min_child_weight': 0.001,
      'min_split_gain': 0.0,
      'n_estimators': 100,
      'n_jobs': None,
      'num_leaves': 1248,
      'objective': None,
      'random_state': 42,
      'reg_alpha': 0.0,
      'reg_lambda': 0.0,
      'subsample': 1.0,
      'subsample_for_bin': 200000,
      'subsample_freq': 0,
      'verbose': -1,
      'n_estimator': 10,
      'min_data_in_leaf': 100,
      'feature_fraction': 1.0,
      'bagging_fraction': 0.6}
```

1.3.6 Stacked model:

```
[]: from mlxtend.regressor import StackingCVRegressor
```

Define component models:

```
[]: directory = 'trainer_checkpoint_data/'
     with open(directory + 'elastic net_checkpoint.json', 'r') as json file:
         data = json.load(json_file)
         elastic_model = ElasticNet(**data['best_params'])
     with open(directory + 'support vector regressor_checkpoint.json', 'r') as ___
      ⇒json_file:
         data = json.load(json_file)
         svr_model = SVR(**data['best_params'])
     with open(directory + 'random forest regressor_checkpoint.json', 'r') as_
      →json_file:
         data = json.load(json_file)
         rfr_model = RandomForestRegressor(**data['best_params'], n_jobs=-1)
     with open(directory + 'xgboost regressor_checkpoint.json', 'r') as json_file:
         data = json.load(json_file)
         xgb_model = XGBRegressor(**data['best_params'])
     with open(directory + 'lightgbm regressor_checkpoint.json', 'r') as json_file:
         data = json.load(json_file)
         lgbmr_model = LGBMRegressor(**data['best_params'])
```

Define blended model:

```
colsample_bynode=None,
                         colsample_bytree=None, device=None,
                         early_stopping_rounds...
                          learning_rate=0.01, max_depth=11,
                          min_data_in_leaf=100,
                          n_estimator=10, num_leaves=1248,
                          random_state=42, verbose=-1),
            SVR(C=1000, gamma=1e-08),
            ElasticNet(alpha=0.01, l1_ratio=1,
                       max_iter=3000, random_state=42),
            RandomForestRegressor(max_depth=1000,
                                  max_features=200,
                                  min_samples_split=25,
                                  n_estimators=1024,
                                  n_{jobs=-1}
                                  random_state=42)),
use_features_in_secondary=True)
```

[]: evaluate_model(stack_gen, 'Stacking model', X_test, y_test, y_logscale=True, user='results/BoW/')

c:\Users\aupho\AppData\Local\Programs\Python\Python311\Lib\sitepackages\lightgbm\basic.py:1218: UserWarning: Converting data to scipy sparse
matrix.

_log_warning("Converting data to scipy sparse matrix.")

Stacking model's evaluation results:

-	Mean squared error:	12.04
-	Root mean squared error:	3.47
-	Mean absolute error:	1.99
-	R2 error:	0.03
-	F1 score:	0.34
-	Precision:	0.46
-	Recall:	0.31
-	Accuracy:	0.31
